

e auropéen des brevets

EP 0 826 631 A1 (11)

(12) FUROPEAN PATENT APPLICATION

(43) Date of publication: 04.03.1996 Bulletin 1998/10 (51) Int CI 5: C01B 39/22, C01B 13/02, B01J 20/18, B01D 53/02

(21) Application number 97306575.8

(22) Date of filing: 28.08.1997

(84) Designated Contracting States: AT BE CHICK ON DIK ES FI FR OB OR IE IT LI LU MC NL PT SE

(30) Priority: 50.06,1996 JP 229676/96 29.11.1996 JP 319819/98 10.03.1997 JP 54710/97

(71) Assigned TORON CORPORATION Shinnanyo-ehi, Yamasuchi-ken (JP)

(72) Inventors: Funakoshi, Helime

Shinnanyo-shi, Yemaguchi (JP) . Shirakura, Yeshinori Shinnanyo-shi, Yamaqushi (JP) Yataunami, Shunauke Shinnanyo-shi, Yamaguchi (JP)

· Yamemoto, Kezueki Tokuyama-ahi, Yamaguchi (JP)

· Ogewa, Nobuhiro lehlicaws-shi, Chiba (JP) · Mort, Takashi

30 John Street

Hikari-shi, Yemanuchi (JP) Harada, Atmushi Shinnenyo-shi, Yemaquohi (JP)

(74) Representative: Kearney, Kayin David Nicholas et al KILDURN & STRODE

London, WC1N 2DD (GB) (54) Heat-resistant low-silica zeolite, and process for production and application thereof

A novel heat-resistant low-silica zeolite, an industrial production process, and uses of the low-sition zeolite are provided. The heat-resistant low-sites zeoite contains Si and Al in a molar ratio of SiO-/Al-Oranging from 1.9 to 2.1, and has sodium and/or potesslum as metal cation, wherein the low-sitics zeolite con-

tains low-sitica faujusite type zeolde at a content of not lower than 88%, and has a thermal decomposition temperature ranging from 870°C to 900°C in the air. The process for producing the heat-registed low-ellics zeoIts comprises mixing a solution containing an aluminate with another solution containing a silicate, allowing the resulting mixture to get, and aging the resulting get, at the lemperature of from 0°C to 60°C, to prepare a sturry having a viscosity ranging from 10 to 10000 cp and confalming amorphous aluminositicate having a specific surface area of not less than 10 mf/g with an SiOy/AlyOs moter ratio ranging from 1 9 to 2.1; and subsequently crystalizing the alumnosilicate. The low-silica zeoite ion-exchanged with lithium or an atkeline earth metal is useful for one percentifico

Description

Background of the Invention:

Field of the Invention:

- The present invention relates to a novel low-silica zeolas of an SIO-/Al-O- molar ratio of 1.9-2.1 having extraordinary high heat resistance. The precent invention relates also to a process for industrial production, and application
- The powel low-ellics zerolin of an SIOvALO-moler ratio of 1.9-2 1 having extraordinary high heat exhauson retains the high heat resistance even after ton-exchange with various ions. This low-effice goodle exhibits extremely high performance, for example, as an actionalism zeolite in separating and concentrating dayugn from an oxygen-nitrugen pag mixture by adsorption, or as a CO2 adsorbent.

18 Description of the Retired Art.

- The low-sition zerolite of an SiOyAlyO₃ molar ratio of 1.9-2.1 (hereinafter referred to as "LSX") is known to exhibit high performance as an adsorbent base material for oxygen production, or an adsorbent for COs gas.
- However, no technique has been established for industrial production of LSX, and the adsorbani prepared by on exchange of an LSX prepared by a known faboratory technique is inferior in heat recistance. Therefore, the LSX has not been practicalized incustrially.
 - The performance of conventional LSX exchanged with fithium for was avaluated in USP 3.140.933, JP-B-6-25527. and USP 5.269.023. The performance of conventional LSXs exchanged with calcium ion is shown in .IP.A-51/254247 IP-A-6-29264 and LISP 5 454 857
 - However, the known LSXs are propared in a small amount over several days by a laboratory swithesis technique. and the production element be practiced industrially. Moreover, the laboratory USXs are not sufficient in heat resistance. The laboratory methods for preparation of LSX of the SiO₂/Al₂O₃ molar ratio of 1.9-2.1 are disclosed by prior art
 - riceumente os halour GB 1,590,929 (corresponding to JP-A-53-8400) discloses a method in which a mixture containing sodium, potaselum, aluminate, and silicate le ervetetized at a temperature lower than 50°C, or is aced at a temperature lower than
 - 50°C and then crystallized at a temperature of 50-100°C. This method requires a time of 50 hours or more substantially for preparation of high-purity LSX, which is not suitable for industrial production. The resulting LSX is not satisfactory In heat resistance.
 - GB 1.580,928 (JP-A-53-8400) investigated in detail the method disclosed by East Germany Palent 43221, However, the resulting LSX also had low water adsorption depactly, and had a low purity. The above patents cover wide ranges of conditions for preparation of LSX. However, the ranges include regions
 - where LSX cannot be recoured. The inventors of GB 1,580,928 (JP-A-59-8400) presented later a activititic paper (Zeokte, 1987, Vol.7, p.451-457) to disclose the switherte of LSX in detail, in that paper, high-purity LSX (97% or higher) was obtained by use of a
 - sealed clastic yearel. However, heat resistance was not improved by this mothed. The synthesis in that document was conducted in a small scale by standing in an over during the steps of from aging to crystalitzation, which is not applicable to industrial production, USP 4.859.217 (corresponding to JP-8-5-25527) discloses a method in which a mixture containing sodium, po-
- taselum, and aluminate is mixed with another mixture containing efficete at a low temperature of 4-12°C, the mixture is allowed to get, and the formed get is aged at 38°C and crystallized at an elevated temperature of 70°C.
- The above patents describe that the delation takes two to three clave, and application of excessive mechanical enemy should be avoided.
 - Even at the time (Application date of USP 4.859,217 (June 30, 1987)), the synthesis of LSX takes long time without application of mechanical energy, namely stiming, and the resulting LSX likelit is less heat-resistent. Alternatively, USP 4,603 040 (corresponding to JP-A-61-222919) discloses preparation of LSX from kaplin as the
- aluming and silice source with eliming. In this method, however, the LSX content is no more than about 50% of all zeolite even after the reaction for 100 hours or more, with 10% or more of A type Zeolite evoluced as a byoroduct, and the adverse effect of stirring is reconfirmed for high-ourity LSX production. Moreover the formed "invariances condeneate" has a particle diameter exceeding 50 µm. Therefore, the resulting LSX, after for exchange, does not give sufficient nitrogen adsorption capacity when it is used for nitrogen adsorption from air in high-purity oxygen production by a pressure-swing adsorption (hereinster reteined to as "PSA"), being not suitable as the base low-size, zeolite for
 - PSA gas expension. The LSX is not useful also for CO₂ gas adsorption for the same reason. At the moment, the low-elics zeolite (LSX) is believed to be producible only by reaction for a long time with stiff-

- stending. No disclosure is found on industrial process for USX, and improvement of the heat resistance of the industrial LSX. The high performence of the advolvent employing the LSX as the base zeeks is confirmed in laboratory only. end has not been realized industrially.
- On the other hand, industrial production of oxygen by the PSA process is practiced in iron production with a blast humace, glass production in a fusion turnace, bleaching, fermentation, and so forth by use of an adsociont in an amount of from tons to several time of tons in one batch. Therefore, the tarm "industrial production" herein means production
- of several tons or more of zeoffic in one batch, not production in several kilograms. Known LSX-based accordents are produced by ion exchange of LSX with lithium cation; alkeline earth metal cation such as calcium cation and strontium estion; or composite cation of lithium pation with another eation such as alkaline
- earth metal cetton. They are produced in a leboratory, and is not heat-recistant (e.g., USP 5,152,813) USP 3,140,923 discloses that faujusts exchanged with lithium ion exhibits high performance in nisogen edecaption.
- higher at a higher lithium ion exchange ratio, and the truspenses of an SiOyAl₂O₃ moler ratio of up to 2.0 are useful therefore. This USP does not mention the heat resistance of the laujasite employed.
- The faujosite exchanged with lithium ion at a higher exchange ratio was availabled further by USP 4,859.217 15 (corresponding to JP-B-5-25527) and USP 5,289,023, and the properties are shown in detail. However, the LSX was prepared by a conventional method in the disclosures, and was less heat-resistant.
- The inventor of the above USP 4,659,217, Chien C. Chao, indicated the low heat resistance of Ethium-exchanged taxiastic in USP 5, 174,979, and reported the improvement of the heat resistance of the faujusite by mixed ion exchange with lithium ion and alkaline earth metal ion. The toujuste obtained by exchange with mixed ions of lithium and an alkaline earth metal, although the heat resistance thereof is improved, has a poor adsorption ability, especially at a low
- temperature, so that the improvement of the heat resistance is of no value. Zeolite adsorbente derived from LSX by exchange with cation of an alkaline earth metal such as calcium and strontium are disclosed in JP-A-61-254247, USP 5,173,462, USP 5,454,887, and so forth. However, they are prepared
 - from conventional laboratory LSX as disclosed by GB 1,590,928 (corresponding to JP-A-53-0400) and other patents, and is not sufficient in heat resistance file the aforementioned filhium-exchanged one.
 - The LSX, having potentiality of high performance, has not been used in industrial gas separation by PSA because of difficulty in industrial LSX production and insufficient heat resistance of the conventional LSX,

Summery of the Invention:

- The present invention intende to provide 8 novel LSX having high heat registerics, and a process for industrial production thereof
- According to an aspect of the present invention, there is provided a heat resistant low-slice zeolite of a molar ratio of SiOy/Al₂O₃ ranging from 1.9 to 2.1 having sodium and/or potestium as a materication, the low-silics zeofite containing low-silice faulasite type zeolite at a content of not lower than 88%, and has a thermal decomposition temperature ranging from 870°C to 900°C in the air.
- According to another expect of the present invention, there is provided a process for producing the above hestresistant low-sitics zerolite, the process comprising mixing a solution containing an aluminate with another solution containing a stricate, allowing the resulting motions to gal, and aging the formed get, at a temperature ranging from 0°C
- to 60°C, to prepare a starry having a viscosity ranging from 10 to 10000 op end containing amorphous eluminositicate having a specific surface area of not loss than 10 mVg with an SiQuAl₂Q₃ molar ratio ranging from 1.9 to 2.1; and subsequently crystallizing the amorphous aluminositicate.
- According to still another aspect of the present invention, there is provided a lithium ion-exchanged low-allicazeolite for gas separation which is derived by exchanging the above heat-resistant low-sities zeolite with lithium ion at a lithium auchange ratio ranging from 75% to 100%
- According to a further aspect of the present invention, there is provided an attelline earth metal ion-exchanged low-silics zeolite for gas separation which is derived by exchanging the above heat-resistant feujaste type tow-silica. zoolite with an alkaline earth metal ion at an alkaline certh metal exchange ratio ranging from 40% to 75%.
- According to e still further aspect of the present invention, there is provided a use of the aforementioned low-slice zeolite for production of high purity oxygen gas, comprising adsorbing nitrogen gas from the air by pressure swing adeorption (PSA).
- According to a still further aspect of the present invention, there is provided a use of the algrementioned low-stice zeolite for separation, recovery, and removal of CO2 gas by adsorbing CO2 from a gas by pressure swing adsorption (PSA)

Bnaf Description of the Drawings:

Fig. 1 shows thermal decomposition charts of low-sites zeoine obtained in Example 1, Example 3, Example 4,

EP 0 826 531 41

- and Comparative Example 1.
- Fig. 2 shows the med decomposition chants of low-size zeolds obtained in Examples 7~8, and Comparative Examples 3~4.

 Fig. 3 shows comparison of the air separation properties (separation coefficients) of low-sizes zeolds of the present
- Fig. 3 shows comparison of the air separation properties (separation coefficients) of tow-effice zeolitis of the private invention with a conventional low-sites zeolitis prepared according to the method of USP 6,174,979.

Detailed Description of the Preferred Embodiment:

After comprehensive studies on technique of industrial production of LSX. We herecome of the present invertical bound that in they may LSX can be obtained by a celebrity (LSX can be obtained by a celebrity of production of mixing the severe marries to control the viscosity of the formed gui, in the ST specific quartice area, and so both, and stimping the gui outling againgted inversement of central production being associated by high patient estimates on the mode LSX varies have a resistance on the production of invertibility to ment for demonst according the actions produced to the production of invertibility to ment for demonst according the actions produced to the production of invertibility to ment for demonst according the actions produced to the production of invertibility to ment for demonst according the actions produced to the production. The production of the production o

has been accomplished based on the above findings.

The process for producing the base zeofile, LSX, of the present invention is described below.

The Invarious of the present invarious studenths process of synthesis of LSEA of SICU_ANCS, make its not in 16.2 and in the control of the present invarious studenth the controllers studenths the controllers studenths to exceed the process. As the security is made to all the controllers students of the process commerting students of extra and students and students and students stud

55 modification.
Further it was bound that the sturry having the above properties can be synthesized is addly by mixing an equipose shall siliciate solution of an SIC_VINs_O+K_O) motile ratio of lower than 1.0 with an appeaus shall sharehase solution of an AIC_VINs_O+K_O) moting ratio of lower than 1.0 in particular, the healt-resident low-shallog agent are soluted to

be producible at the Na₂O(Na₂O_{+x}O) moter retto ranging from 0.5 to 0.75, more preferably from 0.5 to 0.65.

The SiO₂A(₂O_x moter ratio into LSX of the present invention is theoretically 2.0. In consideration of measurement arror in charmost inchmical snakelysis, LSX taking the composition ratio in the range of from 1,9 to 2,1 naturally faller in the acope

of the present invances.

The solution containing an alterninate in the present invantion includes solutions of commercial earlier making in the present invantion includes solutions of commercial earlier making in water, solutions of alumnum hydroxides in a socialism hydroxide, and alumnium component in the alumnium containing solution in the containing solution in solution in the solution in the containing solution in solution in the solution in the solution in solution in the solution is not solved for the solution in solution in the solution is solved to be solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the solution in the solution is solved in the solution in the sol

column is not higher than 30% by weight, perinnelly not higher than 20% by weight, but is not specially intrinsic hineron.

In the above concentration range, the abminded e-containing seletion has a low viscosity, and can be intered residily with
the elibrate containing solution.

The solution containing a visitorie in the present invention includes solutions of commencial 42 accidim missions or the solution of commencial 42 accidim missions or the solution while water, sodium hydroxide or possession.

beauties as the in a counting you work, and interview or the location byte counting and the properties of the properties of the concentration of elicon component in the allicon-centaining solution is preferably not higher than 40% by Weight, more preferably not higher than 50% by weight in terms of SICs, but is not limited thereto. In the above con-

contration range. The elicitate-containing solution has a low Veccetify, and can be mixed readily.

The allowments and instrument of the elicitate solution and the elicitate-containing adultion are witness delicated on the mixed to get the elicitate of elicitate is strowed to get. The temperature of eliciting is in the range of icom O'C to 80°C, prefamility isom 20°C to 80°C.

This temperature of mixed near manners the temperature at the limit of comprehence of the administrate-containing solution with the elicitate-containing readilities. Over of the subministrate containing solution with the elicitate-containing solution with the elicitate-containing solution.

solution with the silicate containing solution. One of the planetate containing solution and the allicate-containing solution with the silicate containing solution and the silicate-containing solution may be as a temperature outside this temperature maps, provided that the temperature of the relative solution with the range of the off to solve of the and off the mindre, and be temperature of the respective ecurs makes is obtained on an off finited.

At the temperature of mixing higher than 60°C, A-hypo zerottle or 9-hypo zerottle or one ceesarily tomed as the

byproduct to render difficult the synthesis of a single phase LSX, whereas at the tempositive regions (non-YC 1097C, LSX can be proceduced, but the tower impressive regions used on exponence conding oppositive service as an enforcement of the single phase of the synthesis of th

containing solution is charged into a reaction vosed, and thereto an atuminate-constraining solution is added, in a sale another exemple, water is thereped from a reaction vested, and therefor an atuminate containing solution and self-precontaining solution are added concurrency. The method of initiality and the order of the mixing are not limited provided that the objects of the present inventions can be achieved.

The molar composition ratios at the end of the mixing are preferably as below:

SIO ₂ /Al ₂ O ₃	1.3-22
(Na ₂ O+K ₂ O)/SiO ₂	2045
	0.5-0.75
H ₂ O/(Na ₂ O+K ₂ O)	10-35

more preferably.

25

Na₂Or(Na₂O+K₂O) 0.5-0.65

Orbitals the schow respective ranges of the most protect state, populates such as Angha abolita, sociation, social Proper section are produced in leng amounts, and the pure to revisit costain of SCA/ALA, of their 1-5.4.1 center on leading that been considered generally that the high-purity LEX will not be formed in the about Nu₂Orbita₂O-K₂O-K₂O-Size compa particulus preferred in the present inventor for exempts, in the indust radius buy, the restall of LEX ormation is lower from IDX's according to the document, Zeolin, 1987, Val.71, e.4.50, p. 8, 4.1. kis his, high-purity LEX cannot be produced by a conventional sill-interface decirings (see Company) to Exemptic 20.

By mixing the source materials in such a manner, amorphicus eluminosticate get le generally formed during or after the mixing of the source materials.

The LEX production process has been considered to compress, associately, the stopp of intoing the source material solutions to prepare more a uniform management colution of the compresents, elevating in being produced in a souling, stopping stirring at the time when period heights, and leaving the solutions standing for aging valence supplying entirely all the time when period heights, and leaving the solution standing life aging the supplying entirely all the service of the solution of the solution of the solution begins, aging in allewed to proceed by satisfacting with the writing by legical process by satisfacting with the writing by legical process by satisfacting with the writing or legical process by satisfacting with the writing or legical process of the solution of the soluti

standing to detain the LBX of the SC/ALCs, most rails of 1.9.2.1. However, state completions unsessignion no consistant or the uniformity of the gal, it has been found by the investors of the present leveration that a rowal high early theat resistant LBX can be produced by berning a uniform starty of the critic registration grade gall below the gall protectly eithigh a certain among for example, by interess starting. This completely different from the traditional concept that the setting saids at the final making slep and the avoidance of excessive mechanical energy wageful from the produced size as improved.

In the present investion, the conditione of mixing and string the source manageals are executedly selected on an interior to determine the condition of the complete form to 1000 cope. All the subject selected principle for a 1000 cop. the always a point splatford and party stoppens in the number vested to impair the unationing of the exist and 1000 cop. the always a point splatford and party stoppens in the number vested to impair the unationing of the exist and principle register that the condition of the condition of the condition of the stoppens of the selection of the condition of the con

methods without initiation.

Subsequently, the formed surry is subjected to sight, The aging lamperature range prolessing and to committee.

Subsequently, the formed surry is subjected to sight, The aging lamperature range prolessing the subsequently, the formed surry is subjected to sight, the aging does not take longly line, and the impurity formation within may occur et a higher temperature can be prevented. Specifically the aging is conducted at a tomperature of the may be aging the subject to the subject t

The formed entirely uniform sturry having the viscosity of 10-10000 op may be agod with stirring. The sturry after reaching the aging temperature may be agod with offring or without stirring since the entire sturry has become uniform stirring since the entire sturry has become uniform

After limited investigation, I was stand by the envelope of a process investion has the freewest of the formed memoplesco althrosticulates operations with a superior standars and on the less than 10 million and make a page, and he standard model investigation and the standard sta

EP 0 825 531 A1

- The yield ratio of LSX is improved by controlling the source material-mixing conditions so as to obtain the smorphose aluminosilicate particle composition approximate to the composition of LSX. In other words, the SiCs/AlsOs moler ratio of the amorphous aluminositicate particle should be in the range of 1,9 to 2,1. Outside this range, Artype
- zerotte or Putype zerolite is produced inestrebly undesirably The inventors of the present invention found that the get of the SiQs/AlgQs motar ratio of 1.9-2.1 having a BET specific surface area of not less than 10 mP/g after the aging can be produced by mixing an aqueous alical sticals.
- solution of the SKO₂/(Na₂O+K₂O) moler ratio of less than 1.0 with an equeous stall aluminate solution of the Al₂O₂/ (Na₂O+K₂O) malar religion less than 1.0, with less adverse effects of vertation of the mixing conditions. By this method, the LSX cen be produced stably without variation in the BET specific surface area or the composition of the emorphous stammosticate, independently of slight wantion in mixing conditions.
- From a mixture of an aqueous alkali ellipate solution of the SiCly (Ne_xO+K_yO) molar ratio of less than 1.0 with an approus sited pluminate solution of the ALO-VINs-OHK-O) moter ratio of less than 1.0, amorphous aluminositicate can be produced which is composed at extremely fine particles, and has a composition of SIO-VAI-O- motar ratio of
- 1.9-2.1 and a BET specific surface area of 10 m²/g or more independently of the temperature and time of the mixing. 10 At the molar ratio of SiCu/INs₂O+K₂O) or Al₂O₂(Na₂O+K₂O) of not less than 1.0, the surface area and the composition of the formed amorphous atuminos ilicate vary greatly, which makes difficult the stable production of fine amor-
- phous aluminositicate of SiOy/Al₂O₃ molar ratio of 1.9-2.1. in the next step, the get efter the aging for the prescribed time is subjected to temperature elevation for crystalli-
- zation In industrial large scale production, the gel may be heated with stirring eince the heat transfer into the interior of
- the gel requires long time. In this case, the temperature of the get is pretentily elevated in a possible shortest time to prevent impurity formation, specifically is not longer than three hours, more preferably not longer than one hour. The method of the temperature elevation of the get is not limited provided that the get can be heated to the crys-
- taffization temperature in a short time. In an Example, the gelation-aging is conducted in a vessel, and crystallization is conducted in another vessel, and the selfs transported from colation-sping vessel to the grystalization vessel by a pump or the like through a heat exchanger for heating.
- The crystallization is preferably conducted in stationary state without stirring to prevent impurity formation. The crystallization temperature is preferably in the range of from about 60°C to about 90°C to shorten the crystallization time and to prevent impurity formation at a higher temperature
- The time for the crystalization depends on the prostalization temperature, and usually cappas from 4 to 12 hours. in the present invention. A longer time is acceptable, When the crystalization time is shorter, the crystalization can be
- incomplete to leave amorphous eluminositicate uncrystalized The LSX of SiO-/Al-O- moler ratio of 1.9-2.1 having availabled as above is collected by filtration, washed, and
- dried. The filtration, washing, and drying can be conducted by conventional methods. The present invention makes practicable the production of the LSX of SKO₂/Al₂O₃ motor ratio of 1.9-2.1 without cooling of the source material mixture and with poing under stirring, making feasible the industrial production of the powel beat-resistant LSX
- The LSX of the present invention has the thermal characteristics and the application fields as described below. The novel LSX of the present invention retains the heat resistance even after exchange with another ion, and the heat resistence is much higher then conventioned LSXs, surprisingly,
- For example, LSX derived by exchanging the LSX as the base zeolite of the present invention with lathium ion and/ or alkaline earth metal ion has extremely high performance in adsorption separation of a more polar gas component in a gas mixture. The exchange ratio of the LBX of the present invention with lithium can be in the range of from 75% to 100%, or
- enably from 75% to 68%. Although the higher tithium exchange ratio gives higher performance, the production cost will be high gwing to expensiveness of fithium. The exchange rate of the LSX with a leatine earth metal cation is preferably ranges from 40% to 78% in the present
- invention The lithium-exchanged LSXs and the alkaline earth metal-exchanged LSXs are novel substances having excellent
- have resultance, and are different from the low-size zentities or LSXs disclosed in USP 3.140.933, USP 4.859.217. USP 5,258,023, USP 3,140,932, USP 5,179,462, USP 5,454,857, and JP-A-61-25424. The ion exchange of the LSX of the present invention with lithium ion or alkaline earth metal ion can be conducted
- by a conventional process without employing a special procedure, although if is completely different from conventional LSXe in thermal characteristics The thermal cheracteristics of the zeofite are evaluated conventionally by differential thermal enalysis (DTA) as
- described in USP 5.174.979 UP-A-7-255094, and so forth. The thermal decomposition temperature in the present invention is defined as the peak top temperature (first thermal decomposition temperature) of the lowest temperature detectable by DTA. By this generally accepted evaluation

method, the novel LSX of the present invention and the low-exchanged products have higher heat resistance than the conventional LSX and low-exchanged product under the state into exchange condition. The thermal decorposition temperature of the LSX of the present invention in in the range of trom 2070 to 300°C, and is higher by at least 5°C, \$60,000°C, and is higher by at least 5°C, \$60,000°C, but the state of conventional LSXs. The treatmet decomposition temperature of the presentance of the presen

generally for 0 then that of conventional LSX. The based decomposition temperature of the ion-exchanged LSX of the persent invention also a higher than that of the corresponding conventional ion-exchanged LSX years (five. The purity of the LSX of the present invention is pretentably 95% or higher, many prefusely 100% (single phase). The purity of LSX is evaluated mainly by X-ray orbital extracture analysis, and is represented by the ratio of the office that present improving the bits of the LSX. The control of the inspurities earls no core checked by decreased or where described present improving the bits of the LSX. The control of the inspurities earls no core checked by decreased or where described.

rethret to the single phase LSX.

The addorted employing the novel LSX of the present invention as the base zookle addorted a retained polar gas from a gain motive, and is useful for expansion and purification of gases, such as production of high-purity oxygen by addoctorion of minings from the sit, and removal of harmful caddon monotonic form a combustion gas. The addorted is especially useful in pressure-employ addoctory for production of the combustion gas. The addorted is especially useful in pressure-employ addoctory of production of the combustion of the com

is capcularly useful in pre-sure-wind pre-sure-wind pre-sure-wind pre-sure-wind pre-sure-wind pre-sure-wind of Co.g. gas.

The present invention is described below in more detail by reference to Examples and Comparative Examples without limiting the invention.

The measurements were conducted as below in Examples and Comparative Examples.

(1) Chemical Constitution

The sample was dissolved by nitric acid-hydrefluoric acid. The chemical constitution was measured with this solution by ICP emission specifichemical analyzer (model: Optims 9000, Perkin Elmar Co.).

(2) Crystal Structure:

The crystal structure was measured by an X-ray diffraction analyzer (model: MXP-3, MecScience Co.).

(3) Moisture Equilibrium Adsorption Capecity;

A powdery sample dried at 100°C was left in a desiccator at a rotative humidity of 80% for 16 hours or longer, end the sample was heated at 900°C for one hour to measure the decrease of the weight.

(4) Viscosity

The viscosity was maissured by 8-type viscometer (menufactured by Tokimokku K.K.) according to JIS-K-7117 (issued on 1997).

(5) BET Specific Surface Area:

The BET specific surface area was insasured by the apparatus, Flowsorb II 2000 (Micromerities Co.).

(6) Thermal Characteristics by DTA-TG:

The thermal characteristics was evaluated by differential thermal analysis, which is generally employed for heat resistance evaluation of zeolitis, with a differential diversel belance, Thermolise, Model §100 (Flagabu Denki KK) in the ori of a temperature selevation rate of 10°C/cmir up to 100°C/c at an aff they size of 50° michigan.

(7) Air Separation Characteristics,

The lon-destrumped LTXX proposed according to the present invention was evaluated to statisticate patients as explain (according to the present invention was evaluated to statisticate properly and super-secreted by the separation conflictest at 0.4 kg/bm², the pressure adopted generally as the adoption present in industrial presents evaluate conflictest at 0.4 kg/bm², the pressure adopted generally as the adoption presents in a facilitation and the pressure evaluated proposed. The supersection conflicted is derived according to the formula; of the facilitation of the supersection conflicted.

Separation coefficient

Nitrogen adsorption volume × Oxygen partial pressure Oxygen adsorption volume × Natrogen partial pressure

(1)

The adsorption pressure conditions for PSA generally employed were decided in consideration of the average of the data shown in JP-A-9-71950, JP-A-9-239204.

Example 1

In a 5 of an alleries silest reseal or reseal, were pieced 523 of equipment accident multimiter selection (16,000, 2000, 4000,

29 sturry was covered, and was placed in a sir dryer topt at 70°C for 8 hours for crystalization. The resulting prystaline matter use weather with pure water sufficiently and was dried at 10°C oversignt.
The obtained powdery crystal was found to be a single-phase faulasis type zeofts by X-ray cerraction enalysis.

The obtained powdery crystal was found to be a single-phase faulastic type zeotte by X-ray diffraction analysis, having a chemical constitution of 0.87Ne₂O.0.93K₂O.A₂O₂O.205(O₂ and exhibiting moisture equilibrium adeoption of 20.4%. The experimental conditions and the results are shown in Table 1.

Example 2

pe.

The experiment was conducted in the same memor as in Exemple 1 sercept that the aging temperature was clead to 50°C, and the aging temperature shorteand of a Nove because of the higher aging impressures. At the end of the aging, the vessority was 240 pp, and the BET people surface serve was 240 pp, and the BET people surface serve was 240 pp, and the BET people surface serve was 240 pp, and the BET people surface serve was 240 pp, and the BET people surface serve was 240 pp. The solution power of the serve of the surface serve was 240 pp. The solution power of the serve that was sometimes to a single-phase surjuinate by paying office on analysis, and the server of t

having a chamical constitution of 0.67Na₂O 0.39K₂O Al₂O₃2.09K₃ and exhibiting moisture equilibrium adsorption of 93.2%. The experimental conditions and the results are shown in Table 1.

Example 3

In a 3-lb are saidness staff instriction vessel, were plotted 80g of colorous scokum initiation activities used in Europea, 1,000 g of were 2,000 g of were 2,

the aging was transported with healths, opsialized, seathed and ched in the same majors as in Example 1.

The obtained powders operative sections to be a single-glesse insighale layer section by X-ray difficusion analysis, having a chemical constitution of 0.61%,0.0.3%(X-V_x-Q_-0.65C_x) and additioning moniture equilibrium advolption of 3.44%. The experimental conditions and the results are shown in 1500.

Example 4

The source material composition was the same as in Example 1. A scale up test was conducted with a reaction vessel of 30 m² (casely states: 10,00%) This source materials were introduced into the reaction vessel with staming. At the end of immidstation of the source materials, the state prespectative was 40°C, and the stary viscosity was 50°C, p. The starry was trather starred confriences for aging at 40°C for 12 hours. After the aging a somple was staten out, and solid matter was collected by filliation, washing and endrogled. The get late is proclif sustained asset at 25 m²/₂ m²/₂.

the moler ratio SIO₂/A₂O₃ of the gel was 2.02. The gel story other the aging was transported through a pipe to a separate 90m9-crystalization vessel, During the slurry transportation, high temperature steam was introduced into the transporting pipe to heal the get sturry. The get sturry temperature in the crystallization vessel was 70°C. The transportation and heating was conducted in 30 minutes. After the completion of transportation and heating, prestatization was allowed to proceed by still-standing without stirring for 8 hours.

The resulting crystalline matter was washed with water, and dried at 100°C. The obtained powdery crystal was found to be composed of a faulashe type zeolite at a purity of 99% by X-rey diffraction analysis, having a chamical constitution of 0.69Na₂O-0.32K₂O-Al₂O₃·2.08iO₂ and exhibiting moisture equilibrium adsorption of 39,1%. The one beign produced 2.4 ions of LSX in about one day. The experimental conditions and the results one shown in Table 1.

Thus the process of the present invention was confirmed to be capable of producing a large amount of high-curity LSX in one batch industrially.

Example 5

- In a 3-liter stainless steel reaction vessel, were placed 453 c of acaseux socium aluminate solution (Na.C: 20 0 w/5, Al₂O₄, 22.5 w/5, Al₂O₄Na₂O: 0.68 (motar ratio)), 923 g of water, 235 g of sectium hydroxide (purity: 99%), and 215 g of polessium hydroxide (Iral-grade reagent, purity 85%). The content in the vesse) was stirred at 90 mm with cooling on an ice water both (about 2°C).
- To this solution, was added 898 g of equicous sodium siticate solution (Na₂O: 3.8 w/%, SiO₄: 12.6 w/%, SiO₄: Na₂O: 3.4 (moler ratio)) and 137 g of water, both being cooled with Ice, in a time of 5 minutes. The solution after the addition of the sodium silicate was transluced. After stirring for further 20 minutes, the temperature of the water bath was elevated to 36°C. When the solution temperature reached 25°C, the solution began to get with white turbidity. At that time the stirring was intenzified to 250 rpm. Although the viscosity of the gal rose as a whole, no local get stagnation
- occurred in the reaction vessel. The viscosity of the gal was 120 cp. The gal was aged at 36°C for 48 hours with continuous slining. The gol clumy after the aging was transported by means of a roller gump with heating through a stainless steel spiral heat exchanger (8 mm diameter) immersed in an oil bath kept at 80°C into a 3 liter stainless steel crystalization vessel. The temperature of the gal at the inlet of the crystalization vessel was 70°C. The transportation of the gal sluny with heating was completed in 20 minutes.
- Then the crystallization vessel containing the gel story was covered, and was placed in a air dryer kept at 70°C. for 16 hours for crystalization. The resulting crystaline matter was washed with pure water sufficiently and was dried at 100°C overnight. The obtained powdery crystal was found to be a single-chang lauriants home zerolite by X-ray df. fraction analysis, having a chemical constitution of 0.76Na,O 0.24K₂O-Al₂O₂-2.05KO₂ and exhibiting moisture equilibrium adsorption of 33.1%. The zaolite in which the entire of the potassium was replaced by sectum exhibited moisture
- equilibrium adsorption of 35,4%. The experimental conditions and the results are shown in Table 1. As shown in this example, the LSX can be produced stably without formation of impurities, even when the aging time or the crystallization are lengthened,

Example 6

- In a 3-litze attainless steel reaction vessel, were placed 453 g of equeous sodium aluminate solution (Na₂C: 20.0 wr%, Al₂O₃: 22.5 wr%, Al₂O₃/No₂O: 0.68 (molar ratio)), 1060 g of water, 295 g of sodium hydroxide (purity: 99%), and 215 g of poinssium hydroxide (first-grade reagen), purity, 65%). The contest in the vascel was stirred at 250 mm at a temperature of 36°C on a water bath. To this solution, was added 839 g of aqueous sodium silicate solution (Na,O 3.8 wt%, SiO_{xt} 12.6 wt%, SiO_y(Na₃O, 3.4 (motor ratio)) in 5 minutes and 50 seconds. Three minutes after the start of
- the addition of the selectic solution, the mixture became jurbed and began to get. During the addition of the sodium allicate solution, the viscosity of the get rose as a whole, but no local get stagnation occurred in the reaction vassel. At the end of the addition, the slurry temperature was 35°C, and the viscosity thereof was 320 cp. The get was aged at 36°C for 24 hours with continuous eliming. After the aging, a sample was taken out from the clurry. The solid matter in the sample was collected by fitration, washed, and analyzed. The specific surface ones was 20 m²/g, and the SiO₂/ Al₂O₃ molar ratio was 1.98. The gal slurry after the aping was transported by means of a roller pump with heating through a stainless steel spiral heat exchanger (8 mm diameter) introcreed in an oil bath kept at 80°C for heating into a 3-liter stainless stool crystalization vessel. The temperature of the gel at the injet of the crystalization vessel was
- 70°C. The transportation of the gol slurry with heating was completed in 20 minutes. Then the crystalization vessel containing the get alony was covered, and was placed in a air diver kept at 70°C for 8 hours for crystallization. The moulting crystalline matter was weshed with pure water sufficiently and wes dried at 100°C overnight. The obtained powdery crystal was found to be a single-phase faulests type zeolite by X-ray diffraction analysis.

having a chemical constitution of 0.76Na₂O-0.24K₂O-Al₂O₂-2.05O₂ and assisting moisture equilibrium adsorption of 33.4%. The experimental conditions and the results are shown in Table 1.

Comparative Example 1

In the minimar resistance in LVP 4-8,09.27 r (Jin A. 4,5827), 03.19 g of an appeacus confum ristance solution (Ni₂C. 30 sets), 50.0; 24.6 sets), 50.0; 24.6 sets, 50.0; 24.6 sets, 50.0; 24.6 sets, 50.0 sets,

In a state of outstand pudding, and became complete sold in 5 minutes.
The pudding-line power was essented in the vessel, and was aged at 39°C for 48 hours in a thermostated driet, and then crystallactified at 70°C for 16 hours. At the end of the sign, a sample was taken out from the situry. The sold matter in the sample was collected by firstline, weathers, and analyzed. The specific outsize area was 60 firthing and the SIGV.
ADA, more in the sign was 20°C the sold matter was introopted. Altertide, charitized, was shed, and detect in the sample.

manuser as is Example 1.

The obligation dipositiony crystal was found to be composed of faujustic type zeotite at a purity of 99% by X-ray differention destylate, believing a chamical constitution of 0.73/ks_Q-0.28%_Q-0.45%_q-0.95%_g and exhibiting measures equilibrium addression of 23.1%, the expendituation conditions and the regulate are shown in Table 1.

20 Comparative Example 2

Scale-up of the process of Comparative Example 1 was fried by use of a jacketed 500-lifer reactor under the serve conditions. The temperature abvastion of the formed get was sted from the get preparation temperature, 5°C, to the

soling temperature, 30°C, by elevating the jacket temperature. Even after 12 hours, the temperature of the interior got portion was not higher than 50°C, athrough the temperature of the upperature was up portion mean the reactor wall was 50°C. After 48 brows, the temperature of the owner portion was not higher than 50°C.

After 48 hours, the scent imperature was either eferorated to 80°C. However, after 8 hours, the temperature at

this centric portion was not higher than 40°C, although the temperature of the persion may the neasons wall have 70°C. The reaction product obtained with such temperature destribution was evaluated by X-ray distribution. The product at the central portion was an available gold of mornipated reaction, and the product on the wall of the exaction vessel was a miditure of the injuries X-ray possible upon a 40°C and offer imprettle with title learning on the invertible tollage the possible. Thus, untitarnity of the reaction system was bound to be assential for pyrithesis of the high-purely LSX, and application of monthand owney by derived is selected in the type coals production.

35 Comparative Example 3

In the menner as described in Zeotte, 1997, vol.7, p.451., the material composition of Example 2 was aged by still standing at 50°C for 24 hour, and was ceptibilized at 100°C for 3 hours by still standing.

At the end of the eight, a sample was taken out from the elary. The sold matter in the sample was collected by

Elization, washed, and analyzed. The specific ourface area was 70 m²/g, and the SiQ/Al₂Q₃ motion ratio was 2.02. The agod matter were transported, heated, crystalized, washed, and direct in the same manner as in Example 1. The detelhed powder or yested was found to be in a mixed phase state composed of a small amount of an LSX

The obtained powdey crystal was found to be in a mixed phase state composed of a small amount of an LSX phase and a large amount of impurities by X-ray diffraction amplies, and to orbits moleture equilibrium adsoption of 19.0%. The apparential conditions and the results are shown in Table 1.

The above method did not produce LSX as described in the Bigraium (Zeofile 1997 Vol.7 P.451).

Examples 7~8 and Comparative Examples 4~5

Producty materials were prepared from the powdrey LSIA of Exemple 4 and Comparative Scarge 15 by exchanging the 15.24 km it imms also asknape sind 45%, for the discious also necknape and or 45%, for powdrey LSIA of a Exemple 4 and Comparative Exemple 5, the ten of Reimmendenaged LSIAs and the calcium-exchanged LSIAs

The previous are there in Table 1, and the lypical thermal decomposition charts are shown in Figs. 1 and 2.

The previous are there in Table 1, and the lypical thermal decomposition temperature higher than that of conven-

Example 9

An LSX observed in Eximple 1 as the base material was exchanged with lithium at an exchange ratio of 90%. Separately, another on-exchanged LSX having hast resolutions was prepared by low-exchanged of econventional LSX is with lithium for anticlother in fillian 60%, and cellular 150% excepting to be method disclosed at USS 5,17,475. The two low-exchanged LBXs were resend for all expension characteristics (reparation coefficient). The results are shown in 15.2.

The ion-exchanged LSX of the prior art had exhibited lower performance, especially at lower temperatures.

Example		8xs	Experimental conditions	condi ti	one				
Ž	Sturry	Sturry		Aging	DO		Crystallization	tion	
	Temper-	Mixing	Temper-	Time	>	Stirring	Tempar-	tine	
	(30,	frantaged	Co	θE	(45)		(C)	Chr.)	
Example	1								
	38	5:50	36	16	320	Stirred	į,	۰	
2	38	5:50	20	8	240	Stirred	2		ور
m	10	5:30	36	16	180	Stirred	02		υ8
•	45	2:00	43	12	300	Stirred	2		26
	2	5:00	36	48	120	Stirred	2	4	63
10	36	5:50	38	24	320	Stirred	20		1 A
žž									
Comparative Example	sample								
	10	2:50	36	48	Charasturable	ouck	02	4	
2	ın	2:50	36	48	Unmeasurable	None	2 6		
	ıs	2100	20	24	Unnessurable	None	100	ņ	
žž									

EP 0 826 631 A1

			,	,	s	p	8		,	s
7 (Table 1 (continued)									
ale l				coperties						1
	Ance	Amorphous Uninosilicate		Poudery crysts	stal	fi	Exchanging	Di.	Therm	4
	Gel BET Surface	NET S.10./	X-ray diffraction	ton Equi	Equilibrius moisture	2	K	8	sition temp. *	6 *

Commonto		1																
20 2. 1.00 (1971) plane 3.1. 1.0 (1971) plane 3.1 (1971) pla					,	•	,				,	8					,	
20 1.00 (2014) 1914 1914 1914 1914 1914 1914 1914 1				•	ı	,	١	,	•		66	,			,	,	,	99
20 1.00 (100 to 100 to			ê	3	2	33	33	24	24		,	ю		:	RZ	28	,	,
14/12 1311 131			8	6	ò	Z	63	20	26		-	33			2	72	,	ri
20 1.98 2.02 2.02 2.02 2.03 2.03 2.03 2.03 2.03	(8)		33.4		7.00	33.4	33.3	33.1	33.4								19.8	
20 1.98 2.02 2.02 2.02 2.03 2.03 2.03 2.03 2.03			Single phase	Single phase	armid out	erud erbure	Single phase	Single phase	Single phase					Starte about	2000	TARE PROBE	arxture arxture	
17/11 1	retter																	
Example 10 Comments of the second of the sec	(10,73)	1	20	24	23	i	9 5	9 6	9				anthe	99	56	5	3	
		Example	-10	7	en	-			,	4		,	Comparative Ex	1	64	e		♥ 10

* Pirat thermal decomposition, center temperature

ED 0 828 021 41

Claims

- A hear-resistant low-sitios zeotite of a motar ratio of SiOyAl₂O₂ ranging from 1.9 to 2.1 having sodium and/or
 potassum as a regular cation, wherein the low-sitios avoids contains low-sitios allocation type zeotites as a contain of
 not lower then 80%, and has a sharmed decomposition temperature ranging from B7VG to SiOYC in air
- The heat-resistant low affice according to claim 1, wherein the low-affice genitive contains the low-affice toutselfe type zooile at a content of not lower than 90%.
- 3. A proof is for producing the hear-resistant low-effor zeotite as set forth in claim 1 or 2, comprising midding a solution containing a set forth in claim 1 or 2, comprising midding a collection collection as a closure, allowing the resulting midding to get, and agree 1000 per producing the collection and the c
 - an SC₂/A₂O₃ molar railo ranging from 7.5 to 2.1; and autrequently operationing the amorphous attentionalisate.

 The process for producing the hair resistant low-since zeroine according to claim 3, wherein an appeare sided sitiose obtained on a SC₂/A(A₂O₃-X₂O) and the racio of lower than 1.0, and an appears of this is distincted arbitrary of the side of the side
- an Augo-yNago-K-co) moler ratio of lower than 1.0 are mixed.

 5. The process for producing the above hear-resistant low-sitics zoottle according to claim 3, wherein the Naso's
- (Na₂O+K₂O) moter ratio of the mixed solution is not less than 0.00, but less than 0.75.

 5. The process for producing the heat-redistant low-stice zeofile according to claim 5, wherein the Ne₂O(Na₂O+K₂O)
- moter rollo is not less than 0.50, but less then 0.65
- A limblum lon-auchanged low-elicia scaller for gas separation, derived by exchanging the sex-resistant tow-elicia scale as forth in claim 1 with limblum ion at a tinium archange ratio ranging from 75% to 100%.
 The influence-exchanged low-sitios resolites for east sessariation according to obtain 7, wherein the filthium exchanges
- ratio ranges from 75% to 89%.

 9. An abiline earth metal for exchanged low-effice zooths for one expansion, derived by exchanging the heat-re-
- An essaine earth metal for-exchanged tow-ence zoots for gas experience, censed by exchanging the head-resistant love-sities zootife as sof forth in claim 1 with an alkaline earth metal for at an elikaline earth metal exchange ratio ranging from 40% to 75%.
- A use of the low-sitios zootile as set forth in any of claims 7 to 9 for production of high purity oxygen gas, comprising adsorbing nitrogen gas from the eir by prosture swing adsorption (PSA).

9))	Europeen Patent Office	EUROPEAN SEARCH REPOR	BT	Application Number EP 97 30 6575
	DOCUMENTS CONSI	DERED TO BE RELEVANT		L
Caregory	Diation of complete will of reference pa	indication where appropriate	Relevant to ofping	CLASSIFICATION OF THE APPLICATION (INCOM)
D.A	GENTER H, KÜML: " low-silica faujasi 2.0)" ZEOLITES, vol. 7, no. 7, 7 J pages 451-457, XPC * the whole docume	te (\$102/A1203 .apprx. lanuary 1987, 00562955	1-7	C01539/22 C01513/02 8013/20/18 8013/53/02
D,A	US 4 859 217 A (CH • the whole docume		1,3,7,8, 10	
١				
D,A	the whole docume		1.3,7,8,	
A	US 5 366 720 A (CA - column 1, line 7	GLIONE ALEX J ET AL)	1,2	
A	FR 2 105 451 A (AL + claim 6 = - page 4, line 1 -	SACE MINES POTASSE)	1,2	TECHNON, FIRES
A	US 4 481 018 A (CO • claims 1,12,13 •	E CHARLES G ET AL)	1,9	C010
A	NO 96 02462 A [ALS * page 7, 11ne 30	EMARLE CORP) - page 8, 15me 25 *	3	
	Тпо ратона околол геден жа			
├	Paradiates	Over the substitute of the banks	لسما	"Larry
_	THE HAGUE	28 November 1997		ondaud. B
7 88	AT SOORY OF CITED DOCUMENT ISSUED, INSTRUMENT IS TO TO COME TO SHAPE A SOON OF THE SOON TO SHAPE A SOON OF THE SOON THE SOON OF THE SO	1 coverages oc	Or september	Asso en er